Study Data Specifications

Revision History

Revision history			
Date	Version	Summary of Changes	
2004-07	1.0	Original version	
2005-03-18	1.1	Addition of specifications for define.xml and SAS XPORT	
		transport files specifications. Changes in document	
		organization.	
2006-03-04	1.2	Update information on annotated ECG waveform data. Delete	
		ecg folder under Specifications for Organizing the Datasets.	
2006-11-27	1.3	Addition of specifications for submitting tumor datasets	
		(tumor.xpt) from rodent carcinogenicity studies.	
2007-08-01	1.4	Addition of hyperlink to information for 3.1.1 datasets	
2009-10-30	1.5	Modified introduction. Additional specifications for submitting	
		data tabulation datasets. Additional specifications for analysis	
		datasets. Revision of maximum file size restrictions. Addition	
		of hyperlink to information for 3.1.2 datasets.	
2010-01-04	1.5.1	Modified directory tree structure for organizing datasets and	
		made minor technical corrections.	

STUDY DATA SPECIFICATIONS

These specifications are for submitting animal and human study datasets in electronic format. Datasets are views of the study data used by reviewers to conduct specific analyses of the study data. They may include both raw and derived data. Because of the unpredictability of the scientific review process, it is impossible to enumerate a priori all datasets needed for review. Prior to the submission, sponsors should discuss with the review division the datasets that should be provided, the data elements that should be included in each dataset, and the organization of the data within the file. Additionally, not all FDA centers have adopted all aspects of these specifications, sponsors are advised to discuss with the reviewing division data needs prior to preparing data for submission.

SAS XPORT TRANSPORT FILE FORMAT

SAS XPORT transport format, also called Version 5 SAS transport format, is an open format published by the SAS Institute. The description of this SAS transport file format is in the public domain. Data can be translated to and from this SAS transport format to other commonly used formats without the use of programs from SAS Institute or any specific vendor.

Version

In SAS, SAS XPORT transport files are created by PROC XCOPY in Version 5 of SAS software and by the XPORT SAS PROC in Version 6 and higher of SAS Software. SAS Transport files processed by the CPORT SAS PROC cannot be processed or archived by the FDA.

Sponsors can find the record layout for SAS XPORT transport files through SAS technical support technical document TS-140. This document and additional information about the SAS Transport file layout can be found on the SAS World Wide Web page at http://www.sas.com/fda-esub.

Transformation of Datasets

SAS XPORT transport files can be converted to various other formats using commercially available off the shelf software.

SAS Transport File Extension

All SAS XPORT transport files should use xpt as the file extension.

Compression of SAS Transport Files

SAS transport files should not be compressed. There should be one dataset per transport file.

Maximum Size of User Supplied Elements

Element	Maximum Length in Characters
Variable Name	8
Variable Descriptive Label	40
Dataset Label	40

CONTENT OF DATASETS AND SIZE OF DATASETS

Each dataset is provided in a single transport file. The maximum size of an individual dataset is dependent on many factors. In general, datasets other than SDTM datasets, should be less than 400 MB; SDTM datasets should not be divided. Datasets divided to meet the maximum size restrictions should contain the same variable presentation so they can be easily concatenated.

Datasets which are divided should be clearly named to aid the reviewer in reconstructing the original dataset, e.g., xxx1, xxx2, xxx3, etc. The files that have been divided and need to be concatenated should be noted in the data definition document. This documentation should identify the range of subject numbers (or other criteria used for division) in the label for each of the divided datasets. For further information on file size limitations for files submitted to CDER, contact cder-edata@fda.hhs.gov.

SPECIFICATIONS FOR SPECIFIC DATASETS AND DOCUMENTATION

Study data are provided using different presentations: Data Tabulation Datasets, Data Listing Datasets, Subject Profiles, and Analysis Datasets.

Data tabulation datasets

Definition

Data tabulations are datasets in which each record is a single observation for a subject.

Specifications

Specifications for the Data Tabulation datasets of human drug product clinical studies¹, are provided by the Study Data Tabulation Model (SDTM) developed by the Submission Data Standard working group of the Clinical Data Interchange Standard Consortium

¹ Here, "drug product" also includes biologic products (submitted as BLAs) that are reviewed in CDER

(CDISC) ². FDA centers and reviewing divisions differ in their use of SDTM data. CDER currently accepts SDTM datasets prepared in accordance with the SDTM implementation guide versions listed in the following table. CBER is currently testing SDTM for clinical studies biologic products and for animal toxicity studies. Individual centers or reviewing divisions may specify the version of SDTM needed for review (see below). Follow the corresponding hyperlink to view the appropriate SDTM and implementation guide.

Each SDTM dataset is provided as a SAS Transport (XPORT) file.

SDTM Model Version	IG Version	Implementation Guide	Support Begins	Support Ends
1.0	3.1	http://www.cdisc.org/models/sds/v3.1/index.	2004-07-01	2010-03-31
		<u>html</u>		
1.1	3.1.1	http://www.cdisc.org/content1605	2007-08-01	
1.2	3.1.2	http://www.cdisc.org/content1055	2009-10-30	

While the SDTM provides a valuable representation that may facilitate review, it does not always provide data structured in a way that supports all analyses needed for review. Sponsors should therefore augment SDTM with analysis data sets as described in the *Analysis datasets* section.

Currently, CDER statisticians perform analyses on the tumor data from each rodent carcinogenicity study, and they need this information provided as an electronic dataset. See Appendix 1 on data elements for the dataset recommended by these statistical reviewers (tumor.xpt). This information will be needed until testing is completed on SDTM for animal toxicity studies.

Data Listings

Definition

Data listings are datasets in which each record is a series of observations collected for each subject during a study or for each subject for each visit during the study organized by domain.

Specifications

Each dataset is provided as a SAS Transport (XPORT) file. Currently, there are no further specifications for organizing data listing datasets.

² CDISC, <u>www.cdisc.org</u>, is an open, multidisciplinary, not-for-profit organization committed to the development of worldwide industry standards to support the electronic acquisition, exchange, submission and archiving of clinical trials data and metadata for medical and biopharmaceutical product development.

Subject profiles

Definition

Subject profiles are displays of study data of various modalities collected for an individual subject and organized by time.

Specifications

Each individual subject's complete subject profile is in a single PDF file. Including the subject ID in the file name will help identify the file. Alternatively, all subject profiles for an entire study may be in one file if the size of each individual subject profile is small and there are not a large number of subject profiles needed for the study. If a sponsor does the latter, the PDF file should be bookmarked using the subject's ID. Including the study number in the file name will help identify the file.

Analysis datasets

Definition

Analysis datasets are datasets created to support results presented in study reports, the ISS and the ISE and to support other analyses that enable a thorough regulatory review. Analysis datasets contain both raw and derived data.

Specifications

Each dataset is provided as a SAS Transport (XPORT) file. Prior to submission, sponsors should contact the appropriate center's reviewing division to determine the division's analysis dataset needs. CDISC/ADaM standards for analysis datasets (http://www.cdisc.org/adam) may be used if acceptable to the review division.

Any requested programs (scripts) generated by an analysis tool should be provided as ASCII text files and should include sufficient documentation to allow a reviewer to understand the submitted programs. If the programs created by the analysis tool use a file extension for ASCII text files other than .txt, the file name should include the native file extension generated by the analysis tool for ASCII text program files, e.g. myRcode.r, mySAScode.sas, etc. If the analysis tool does not save programs in ASCII format, a PDF rendition of the program file should be provided in addition to the program file.

General Considerations for Analysis Datasets

For an individual study, all dataset names and dataset labels should be unique
across the analysis and raw datasets submitted for this individual study. The
internal name for an analysis dataset should be the same as the name shown in
the data definition file.

- Each analysis dataset should be described by an internal label which is shown in the data definition file. This label should clearly describe the contents of the dataset. For example, the label for an efficacy dataset might be "TIME TO RELAPSE (EFFICACY)". At least one analysis dataset should be labeled in the data definition file as containing the primary efficacy data.
- The key variables (subject identifier and visit for datasets with multiple records per subject) should appear first in the datasets. Each subject should be identified by a single, unique subject identifier within an entire application (including tabulation, listing and analysis datasets). Subjects enrolled in a primary study and then followed into an extension study should retain their unique identifier from the primary study.
- When a dataset contains multiple records per subject, a variable for relative day
 of measurement or event and variables for visit should be included. In addition to
 a protocol-scheduled visit variable, include at least two timing variables; a
 character variable describing the visit (e.g. WEEK 8) and a corresponding
 numeric variable (e.g. 8). These two variables are measures of time from
 randomization.
- For unscheduled visits or measurements, numbers are often assigned values between two protocol-scheduled visits. These numbers should be distinct from other visit numbers but retain the chronological order (e.g. two unscheduled visits between visit 3 and visit 4 might be 3.1 and 3.2). The character form of the visit identifier may be UNSCHEDULED or a similar term.
- Core variables should be listed after the key variables and included on each analysis dataset. Core variables include study/protocol, center/site, country, treatment assignment, sex, age, race, analysis population flags (e.g. ITT, safety) and other important baseline demographic variables.
- Variable names and codes should be consistent across studies and where
 feasible, the NCI CDISC Vocabulary should be used. For example, if glucose is
 collected in a number of studies, use the CDISC Submission Value "GLUC" for
 the laboratory test code in all of the studies. The NCI CDISC terminology is
 available at http://evs.nci.nih.gov/ftp1/CDISC/SDTM/. Additional information is
 also available at
 http://www.cancer.gov/cancertopics/terminologyresources/page6.
- The format of variables for similar types of data should be consistent within and across studies. For example, all variables that include calendar dates (e.g. birth date, screening visit date, randomization date, date of death) should use the same format for representing the date.
- For textual data that have been mapped to numeric codes, provide two variables, one with text and one with numeric codes.

 Dates should be formatted as numeric in the analysis datasets even if dates are in ISO8601 or another character format in the raw data. This formatting will facilitate the calculation of duration.

SPECIFICATIONS FOR DATASETS DOCUMENTATION

Dataset documentation includes data definitions and annotated case report forms.

Data definition file

Definition

The data definition file describes the format and content of the submitted datasets.

Specifications

The specification for the data definitions for datasets provided using the CDISC SDTM is included in the Case Report Tabulation Data Definition Specification (define.xml) developed by the CDISC define.xml Team. The latest release of the Case Report Tabulation Data Definition Specification is available from the CDISC web site (http://www.cdisc.org/models/def/v1.0/index.html). Include a reference to the style sheet as defined in the specification and place the corresponding style sheet in the same folder as the define.xml file.

For datasets not prepared using the CDISC SDTM specifications, consult Appendix 2 for information concerning the preparation of a define.pdf data definition file.

Annotated case report form

Definition

An annotated case report form (CRF) is a blank CRF with annotations that document the location of the data with the corresponding names of the datasets and the names of those variables included in the submitted datasets.

Specifications

The annotated CRF is a blank CRF that includes treatment assignment forms and maps each item on the CRF to the corresponding variables in the database. The annotated CRF should provide the variable names and coding for each CRF item included in the data tabulation datasets. All of the pages and each item in the CRF should be included. The sponsor should write *not entered in database* in all items where this applies. The annotated CRF should be provided as a PDF file. Name the file *blankcrf.pdf*.

SPECIFICATIONS FOR OTHER TYPES OF STUDY DATA Annotated ECG waveform data

Definition

Annotated ECG waveform data are raw voltage-versus-time data comprising the electrocardiogram recording, to which have been attached the identification of various intervals or other features.

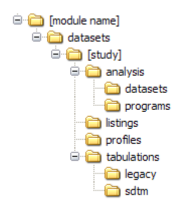
Specifications

See the HL7 normative standard for creating the annotated ECG waveform data files. This information may be found on the HL7 web site www.hl7.org. More information may be found at:

http://www.fda.gov/Drugs/DevelopmentApprovalProcess/FormsSubmissionRequirement s/ElectronicSubmissions/ucm085324.htm#ECG.

SPECIFICATIONS FOR ORGANIZING THE DATASETS

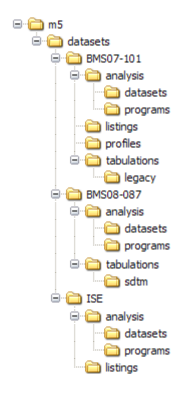
The specifications for organizing study datasets and their associated files in folders are summarized in the following figure and accompanying table. No additional subfolders are needed; unused folders do not need to be supplied.



Folder	Description
[module name]	Name the folder according to the CTD module which the datasets
	apply to. Use m4 for non-clinical data and m5 for clinical data.
datasets	The folder under which all of the study data being submitted for the
	module specified is organized.
[study id]	Name the folder according to the study identifier or analysis performed
	for which the data is being supplied, e.g., BMS09-101, ISS, ISE.
analysis	The folder under which analysis datasets and programs will be
	organized
datasets	The sub-folder in which the analysis datasets are organized
programs	The sub-folder in which the analysis programs are organized

Folder	Description	
listings	The folder in which miscellaneous datasets that don't qualify as	
	analysis, profile, or tabulation datasets are organized	
profiles	The folder in which patient profiles are organized.	
tabulations	The folder under which tabulation datasets will be organized.	
	Note: The tabulations datasets are to be placed in specific folders based on their format.	
legacy	The sub-folder in which tabulations not formatted according an identified standard format are organized, e.g., non-SDTM datasets.	
sdtm	The sub-folder in which tabulations formatted according to the SDTM	
	standard are organized.	

The following example shows the folder structure for a submission containing 2 individual clinical studies and an ISE. Legacy tabulations have been submitted for BMS07-101. SDTM tabulations have been submitted for BMS08-087.



APPENDIX 1

Tumor Dataset For Statistical Analysis ^{1,2} (tumor.xpt)					
Variable	Label	Тур	Codes	Comments	
STUDYNUM	Study number	char		3	
ANIMLNUM	Animal number	char		1,3	
SPECIES	Animal species	char	M=mouse R=rat		
SEX	Sex	char	M=male F=female		
DOSEGP	Dose group	num	Use 0, 1, 2, 3,4, in ascending order from control. Provide the		
DTHSACTM	Time in days to death or sacrifice	num	dosing for each group.		
DTHSACST	Death or sacrifice status	num	1 = Natural death or moribund sacrifice 2 = Terminal sacrifice 3 = Planned intermittent sacrifice 4= Accidental death		
ANIMLEXM	Animal microscopic examination code	num	0= No tissues were examined 1 = At least one tissue was examined		
TUMORCOD	Tumor type code	char		3,4	
TUMORNAM	Tumor name	char		3,4	
ORGANCOD	Organ/tissue code	char		3,5	
ORGANNAM	Organ/tissue name	char		3,5	
DETECTTM	Time in days of detection of tumor	num			
MALIGNST	Malignancy status	num	1 = Malignant 2= Benign 3 = Undetermined	4	
DEATHCAU	Cause of death	num	1 = Tumor caused death 2= Tumor did not cause death 3 = Undetermined	4	
ORGANEXM	Organ/Tissue microscopic examination code	num	1 = Organ/Tissue was examined and was usable 2= Organ/Tissue was examined but was not usable (e.g., autolyzed tissue) 3 = Organ/Tissue was not examined		

Each animal in the study should have at least one record even if it does not have a tumor.

- Additional variables, as appropriate, can be added to the bottom of this dataset.
- ANIMLNUM limit to no more than 12 characters; ORGANCOD and TUMORCOD limited to no more than 8 characters; ORGAN and TUMOR should be as concise as possible.
- A missing value should be given for the variable MALIGNST, DEATHCAU, TUMOR and TUMORCOD when the organ is unusable or not examined.
- ^{4.} Do not include a record for an organ that was useable and no tumor was found on examination. A record should be included for organs with a tumor, organs found unusable, and organs not examined.

APPENDIX 2

Sponsors should include a define.pdf to describe the datasets for each study, specific data analysis (e.g., population PK), and integrated summaries. For the datasets to be useable, the definitions of the variables should be provided. Sponsors should document all of the variables in the datasets in data definition tables. There should be one set of data definition tables for each study, specific data analysis (e.g., population PK) and integrated summary. The first table should include a listing of all datasets provided for the study with a description of the dataset and the location of the dataset file. Provide a hypertext link from the description of the dataset to the appropriate data definition table. Provide a hypertext link from the location listing of the file to the SAS transport file. The reviewer can use the first hypertext link to view the data definition table and the second to open the SAS transport dataset file. Sponsors should also provide a link to the appropriate annotated case report form file (blankcrf.pdf).

In the following table, the dataset for AE is described as adverse events, and the dataset file is located in listings folder for study 1234

Datasets for Study 1234				
Dataset	Description of Dataset	Location		
AE	Adverse Events	m5/datasets/study1234/listings/ae.xpt		

Subsequent pages should contain a table for each dataset that includes an organized listing of all variable names used in the dataset, a descriptive variable label, data types, codes (and decodes), and comments. The comments field is for further description of the variables. For derived variables, the method for calculating the variable should be included in the comments field. For raw variables, the location of the variable on the annotated CRF should be provided as well as the CRF field name if different from the variable name in the dataset. Providing a hypertext link from each raw data variable in the data definition table to the appropriate location of the blankcrf.pdf also helps the review process. An example of part of a data definition table for the demographics dataset for study 1234 is provided below.

Study 1234 – Demographics Dataset Variables				
Variable	Label	Type	Codes	Comments ¹
USUBJID	Unique subject ID number	char		Demographics page 3
SEX	Sex of subject	char	f = female m = male	Demographics page 3
BDATE	Birth date	date		Demographics page 3
DUR	Duration of Treatment	num		Derived STOP DATE – START DATE
TRT	Assigned treatment group	num	0= placebo 5= 5mg/day	

¹Use footnotes for longer comments

The data definition tables should be provided as a single PDF file named *define.pdf* and placed in the appropriate study, specific analysis type or integrated summary folder in the datasets folder. The Title portion of the Document Information field of each data definition file should include the appropriate study report number, specific analysis type or integrated summary name and *data definitions*. For example, the data definition file for study 2001 would be identified as: *study 2001, data definitions*. This file is considered part of the comprehensive table of contents.